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PCT/FR00/018	365	June 30, 2000	July 1st, 1999		
TITLE OF INVEN					
		G ORGANOZINC DERIVA	ATIVES 'SIS WITH COBALT SALTS		
APPLICANT (S) F			315 WITH COBALT SALTS		
		INNE GOSMINI AND YOL	ANDE ROLLIN		
Applicant herewith sul			D/EO/US) the following items and other		
information: 1 ⊠ This is a FIRST	submission of ite	ms concerning a filing under 35 U S	C 371		
2 This is a SECC	OND or SUBSEQU	IENT submission of items concerning	g a filing under 35 U S C 371		
examination u	ntil the expiration	of the applicable time limit set in 35	J S C 371) (f) at any time rather than delay U S C 371 (b) and PCT Articles 22 and 39(1)		
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		cation as filed (35 U S C 371(c)(2)) (required only if not transmitted by the	he International Bureau)		
b. 🔯 has b	een transmitted by	y the International Bureau.	•		
		pplication was filed in the United Sta Application into English (35 U S C 3			
_		h Report (PCT/ISA/210)	. (-)(-)/-		
Amendments to the claims of the International Application under PCT Article 19 (35 U S C 371(c)(2)).					
a. are transmitted herewith (required only if not transmitted by the International					
		y the International Bureau	h amendments has NOT expired		
c. ☐ have not been made, however the time limit for making such amendments has NOT expired. d. ☑ have not been made and will not be made.					
9	9 A translation of the amendments to the claims under PCT Article 19 (35 U S C 371(c)(3)).				
10 ⊠ An oath or de	eclaration of the in	ventor(s) (35 U S C 371(c)(4)). (sig	ned)		
11 🛛 A copy of the	International Prel	iminary Examination Report (PCT/IF	PEA/409).		
12 A translation o		ne International Preliminary Examina	ation Report under PCT Article 36		
Items 13 to 18 below	w concern docum	nent(s) or information included: ment under 37 CFR 1 97 and 1 98			
14 🛛 An assignmen	14 ⊠ An assignment document for recording. A separate cover sheet in compliance with 37 CFR 3 28 and 3 31 is				
included 15 🛛 A FIRST prelin					
16 A SECOND	16 A SECOND or SUBSEQUENT preliminary amendment				
17 A substitute specification.					
18 ☐ A change of p	18				
19 🛚 Certificate of	Mailing by Expres	s Mail			
	6, 308 and 33	2, PCT/ISA/ 220 & 210 (Engl h & French Version). PCT/	iish & French Version), FR00/01865 as published		
Page 1 of 2	, ,	C/Data/RN99079 KMV.Trans.doc			

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a A check in the amount of \$ to cover the above fees is enclosed b Please charge my Deposit Account No. 18-1171 in the amount of \$1,010.00 to cover the above fees. c The Commissioner is hereby authorized to charge any additional fees which may be required, or credit any overpayment to Deposit Account No. 18-1171 A duplicate copy of this sheet is enclosed. NOTE: Where an appropriate time limit under 37 CFR 1,494 or 1,495 has not been met, a petition to revive (37 CFR 1.137 (a) or (b)) must be filed and granted to restore the application to pengling status, SEND ALL CORRESPONDENCE TO: John A. SHEDDEN RAME John A. SHEDEN NAME John A. SHEDEN NAME 25,644						
Cranbury, NJ 08512						
December 20 ,2	2001	REGI	STRATION NUMBER	₹		

531 Rec'd PCT/PTC 2 0 DEC 2001 Case RN99079

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In the Application of: Jacques PERICHON, Corinne GOSMINI and Yolande ROLLIN

National Phase of PCT/FR00/01865

International Filing Date: June 30 2000

Serial No: To be assigned Filing Date: To be assigned Art Unit: N/A

Examiner: N/A

For: PROCESS FOR PREPARING ORGANOZINCIC DERIVATIVES ELECTROCHEMICALLY, COMBINED WITH CATALYSIS WITH COBALT SALTS.

Commissioner for Patents Washington, D.C. 20231

PRELIMINARY AMENDMENT

Dear Sir:

Prior to calculation of filing fee, please enter the following amendment in the

specification and claims:

In the Specification:

Page 1, just after the title, please add the new following paragraph:

This application is an application under 35 U.S.C. Section 371 of International

Application Number PCT/FR00/01865 filed on June 30, 2000.

In the Claims:

Please cancel claims 1-10, and replace them with the following new claims 11-23.

11. (New) A process of electrolytic synthesis of organozine compounds, comprising the use of a catalytically amount of cobalt.

12. (New) The process according to claim 11, wherein, the organozinc compound is an arvl or vinvl organozine compound.

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- 13. (New) The process according to claim 11, wherein cobalt is present in an electrolyte in oxidation state II.
- 14. (New) The process according to claim 11, wherein cobalt is present in a coordinated form.
- 15. (New) The process according to claim 14, wherein cobalt is coordinated with a solutant or solvent compound that has a high donor number.
- 16. (New) The process according to claim 15, wherein the solutant or solvent compound comprises an atom having a high donor number, selected from the group consisting of atoms of the nitrogen column.
- 17. (New) The process according to claim 11, wherein cobalt is coordinated with a specific ligand.
- 18. (New) The process according to claim 17, wherein the ligand comprises a function selected from the group consisting of pyridine, nitrile, phosphine, stibine and imine functions.
- (New) A composition for electrolytic use, comprising a cobalt salt, a zinc salt, a solvent and a cobalt ligand.
- 20. (New) A process for the electrolytic synthesis of an organozine compound, comprising the step of subjecting to an electrolysis on an inert cathode a composition comprising a cobalt salt, a zinc salt, a solvent, a cobalt ligand, and an organic halide.
- 21. (New) A process according to claim 20, wherein the organozine compound is an aromatic or vinyl organozine compound.
- 22. (New) An aromatic organozine compound comprising:

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- an sp2 carbon atom,
- at least one aniline group not more than monosubstituted,
- an SO₂ group, and
- a zinc-bearing function,

wherein the aniline group, the SO_2 group, and the zinc-bearing function are bounded to the sp2 carbon atom.

23. (New) An aromatic organo zinc compound according to claim 22, wherein the sp2 carbon atom is comprised in an aromatic group.

REMARKS

The preliminary amendments are filed to comply with the claims structure and wording according to the United States Patent law. It is asserted that these amendments do not add new matter. Support for these amendments can be found in the specification and claims as originally filed.

Entry of these amendments is respectfully requested.

Respectfully submitted,

December 20, 2001

Rhodia Inc. 259 Prospect Plains Road CN7500 Cranbury, NJ 08512

F/RN99079 Prelim Amend.doc

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DOCKET NO. RN99079

PROCESS FOR PREPARING ORGANOZINC DERIVATIVES
ELECTROCHEMICALLY, COMBINED WITH CATALYSIS WITH COBALT
SALTS

1

The present invention relates to a novel

5 process for synthesizing aryl organozinc derivatives. The invention relates more particularly to the electrolytic synthesis of aryl organozinc derivatives, in the catalytic presence of the element cobalt.

The reactivity of organozinc reagents,

10 especially aryl organozinc reagents, shows many
specificities which would make them particularly
advantageous in many organic synthesis operations.
However, they are difficult to obtain and are often
prepared from organometallic reagents made with more

15 electronegative metals, that is to say more reductive
metals.

In addition, most of the techniques require the use of highly aprotic and especially very dry media.

- In particular, reactions for the electrolytic synthesis of organozinc reagents present the risk of two unwanted reactions: firstly, the reduction reaction to give a hydrogenated derivative, and secondly, a coupling reaction (formation of biaryl).
- 25 A certain number of tests have been conducted in an attempt to perform this synthesis electrolytically. The tests that were the most

conclusive were performed by some of the authors of the present invention.

Mention may be made more particularly, firstly, of the general textbook "Organozinc reagent, a 5 practical approach" (Paul KNOCHEL and Philip JONES Editors, Oxford University Press, December 1998). More particularly, a synthetic route is described therein in chapter 8 by S. SIBILLE, V. RATOVELOMANANA and J. PERICHON (see also Journal of Chemical Society 10 Chemical Communications, 1992, 283-284) and the article by C. GOSMINI, J.Y. NEDELEC and J. PERICHON (Tetrahedron Letters, 1997, 38, 1941-1942).

In these articles, the only route that is described therein is the use of very specific nickel 15 complexes as electrolytic synthesis catalysts, in a limited number of media. However, the use of these nickel complexes, although constituting an important innovation, does not generally make it possible to achieve high yields relative to the haloaryl substrate.

Accordingly, one of the aims of the present invention is to provide a process for obtaining organozinc derivatives in good yields, both in terms of reaction yields (RY) and degrees of conversion (DC). In other words, one of the aims of the present invention 25 is to provide a technique which allows the conversion of the substrate with good selectivity (CY).

Another aim of the present invention is to provide a technique for reducing the reduction and coupling reactions.

Another aim of the present invention is to provide a route that is capable of catalyzing the electrolytic synthesis reaction of aryl organozinc reagents, starting with the corresponding halo derivatives.

These aims and others, which will emerge

10 hereinbelow, are achieved by means of using cobalt as a
catalyst in the electrolytic synthesis of arylzinc
compounds.

According to the present invention, it has been shown that cobalt can be introduced especially

15 into the electrolyte in oxidation state II. Admittedly, cobalt can also be introduced in the form of cobalt III, but, since the medium is a reductive medium, this form will have a tendency to disappear very rapidly to be converted into various species, and

20 especially into cobalt II. The state and form of the catalytically active cobalt have not been completely elucidated.

According to one preferred embodiment of the present invention, it is desirable to use cobalt in the 25 presence of at least one of its ligands.

The coordination of the cobalt is advantageously performed with compounds (solvents or solvating agents) that have a high donor number. More

specifically, it may be pointed out that it is preferable that the donor number D of these solvents should be greater than or equal to 10, preferably less than or equal to 30, and advantageously between 20 and 30, the limits being included. The said donor number corresponds to the ΔH (variation in enthalpy), expressed in kilocalories, of the combination of said aprotic polar solvent or of said ligand, with antimony pentachloride. This is described more specifically in the book by Christian REINHARDT: "Solvents and Solvent Effects in Organic Chemistry - VCH, page 19, 1988". Said page gives a definition of the donor number.

It has been shown, in the course of the study which led to the present invention, that very good

15 results are obtained when the atom coordinating to the cobalt is an atom from the nitrogen column, and advantageously nitrogen. In this case, it is preferable that the ligand atom should not bear an electric charge.

When a specific coordinating agent is used, which does not act as solvent, pyridine, nitrile, phosphine, stibine and imine, or even oxime, functions or groups may be mentioned.

When unidentate (or monodentate) ligands are
25 used, it is desirable to use in the electrolyte a molar
ratio between the ligand(s) and the cobalt which is
high ([lig]/[Co] of about 10 and advantageously ≥ about

100); there is usually no upper limit since the ligands may serve as solvent.

When bidentate or multidentate ligands are used, it is possible to decrease the lower limit to 5 ratios at least equal to 2, advantageously to 4 and preferably to 6, but more preferably to 8.

To be efficient, it is desirable that the cobalt should be present at a minimum concentration at least equal to 10⁻³ M. To be economical, it is preferable that the cobalt should not be too concentrated; thus, it is preferred that the cobalt content should be not more than 0.2 M.

The reaction medium advantageously comprises a solvent; this solvent should be polar enough to

15 dissolve the metals or, more exactly, the metal salts used, and it should be lipophilic enough to at least partially dissolve the substrates from which it is desired to form the organozinc reagent.

It is preferable to use solvents that are

20 sufficiently low in acidity (it is desirable that their
pKa should be at least equal to 16, advantageously to
20 and preferably to 25), so that the reactions with
hydrogen are as limited as possible. Thus, primary
alcohols are too acidic to give very good results.

More specifically, the solvents that will be preferred are "polar aprotic" solvents such as, for example, alone or as a mixture:

- purely oxygenated solvents, in particular ethers, preferably polyethers such as 1,2-dimethoxyethane or cyclic ethers such as THF or dioxane;
- amides or ureas (DMF, N-methyl-2-pyrrolidone,
 imidazolidone, tetramethylurea,
 dimethoxypropyleneurea, etc.);
 - sulfones (for example sulfolane) or sulfoxides (such as DMSO); and
- provided that they are liquid under the operating conditions, nitrogenous derivatives, nitrogenous heterocycles, especially pyridine, and compounds containing a nitrile function (for those that are preferred, see below); and
- provided that they are liquid under the operating conditions, complexing agents (crown ether, HMPT, tris(3,6-dioxaheptyl)amine (TDA-1)), which improve the correct functioning of the reaction by increasing the conductivity, increasing the reactivity of the anion and preventing the deposition of metal at the

Without this explanation being limiting, it would appear that these advantageous phenomena are correlated with the ability to complex the metal cations or as a mixture.

25 As has been mentioned previously, the solvents used may themselves act as complexing agents or ligands. They may especially, and advantageously, contain one or more of the coordination functions mentioned above.

The solvent may be a mixture of an apolar solvent and a polar solvent as defined above by the 5 donor number.

To make the products easier to separate from the reaction media, it is preferable that said solvent should have a boiling point that is substantially different from the compound to be synthesized and from 10 the starting compound.

To facilitate the reaction and to improve the conductivity of the medium, saline electrolytes, occasionally referred to as base salts, optionally modified by the presence of complexing agents, are

15 generally used. These electrolytes are chosen so as not to disrupt the reactions at the anode and the cathode.

According to one of the preferred embodiments of the present invention, an excess of zinc cation, relative to the stoichiometrically required amount, may 20 be used as base salt, advantageously in the form of a fully dissociated salt (in general corresponding to an acid whose pKa is at most equal to 3, advantageously to 2, preferably to 1 and more preferably to zero).

When a soluble anode is used, the electrolyte
25 may be chosen so as to have as cations those
corresponding to the metals of the anode.

The electrolyte may be chosen so as to have as cations metals of high transporting power such as

divalent and advantageously trivalent metals, of the type such as aluminum, on condition that this does not disrupt the base reaction.

As metals used in the base salts, it is 5 desirable to use those that have, besides the stage 0, only one stable oxidation state.

The electrolyte may be chosen such that these cations are directly soluble in the reaction medium.

Thus, when the medium is sparingly polar, rather than

making the metal cations soluble by means of adjuvants, it may be advantageous to use "oniums" that are stable in the electrical inactivity range.

The term "onium" means positively charged organic compounds, the name attributed to them by the 15 nomenclature comprising an "onium" affix, generally a suffix (such as sulfonium [trisubstituted sulfur], phosphonium [tetrasubstituted phosphorus], ammonium [tetrasubstituted nitrogen]). The ones most frequently used are tetraalkylammoniums; the alkyl groups taken in their etymological sense generally contain from 1 to 12 carbon atoms and preferably from 1 to 4 carbon atoms. Phase-transfer agents may also be used.

The anions may be any common anions for inert electrolytes, but they are preferably chosen either 25 from those released by the reaction, essentially halides, or, for example, from complex anions such as perfluoro bis-sulfonimides, BF₄, PF₆ or ClO₄. As a guide, it should be pointed out that DMF, used with, as

a base salt, tetrabutylammonium tetrafluoroborate at a concentration of 0.01 M, gives good results.

Another aim of the present invention is to provide a medium that may be used to perform the

5 electrolysis and to give organozinc reagents. This aim has been achieved by means of a composition comprising at least:

- one cobalt salt,
- one zinc salt,
- 10 one conductive solvent, or a solvent that is made conductive, and
 - one cobalt ligand.

The solvent and the cobalt ligand may be one and the same species, and even a single compound when 15 the solvent is a single compound.

The cobalt content is advantageously between 2×10^{-1} and 10^{-1} M and preferably between 5×10^{-3} and 5×10^{-2} M (closed interval, that is to say limits included).

Not taking into account the organozinc reagents formed, the zinc content is advantageously between 0.05 M and the solubility limit in the medium. When a soluble zinc anode is not used, it may be envisaged that a solid phase consisting of zinc salt(s) is present.

Said composition, when it is used to prepare organozinc reagents, also comprises an aryl halide whose preferred chemical characteristics will be given

later. This aryl halide is advantageously present in a concentration of from 0.1 to 1 M.

It is desirable that the molar ratio
(dissolved species, needless to say not taking into

5 account the organozinc reagents formed) of zinc to
cobalt is between 100 and 1 and preferably between 10
and 2 (closed interval, that is to say limits
included).

It is also recommended that the molar ratio

(needless to say dissolved species, not taking into account the organozinc reagents formed) of zinc to aryl halide should be between 0.05 and 4 and preferably between 0.01 and 2 (closed interval, that is to say including the limits). The lowest values correspond to the case in which a soluble zinc anode is used.

According to one advantageous embodiment of the invention, the intensity and area of the reactive electrode, more exactly of the electrode at which the reaction takes place, are chosen such that the current density j is between 5 and 5 × 10² A/m² and preferably between 20 and 200 A/m² (closed interval, that is to say including the limits).

By routine tests, a person skilled in the art can determine the reduction potential of cobalt in the 25 reaction medium and that of the aryl halide. Once this determination is done, it will preferably be between the reduction potential of cobalt and that of the aryl halide.

The substrates which may be converted into organozinc reagents by the present invention represent a wide range of compounds. The halides are generally halides corresponding to relatively heavy halogens, 5 that is to say halogens heavier than fluorine.

It may also be mentioned as an indication that, when the halogen is linked to an electron-poor aromatic nucleus, it is preferable to use bromines or chlorines as halogen, chlorines being reserved for the nuclei that are particularly electron-poor. If the condition is satisfied by six-membered heterocycles, in the case of homocyclic aryls, to use a chloride, it is preferable that the sum of the Hammett constants σ_p of the substituents (not taking the leaving halide into account) should be at least equal to 0.40 and preferably to 0.50. On the other hand, the nuclei that are particularly electron-rich may use iodine as halide.

For further details regarding the Hammett

20 constants, reference may be made to the 3rd edition of
the textbook written by Prof. Jerry March "Advanced

Organic Chemistry" (pages 242 to 250) and published by
John Wiley & Sons.

The five-membered heterocycles comprising a chalcogen as hetero atom (such as furan and thiophene) have a large capacity to be converted into zinc reagents, show separate reactivity, and are always readily converted into zinc reagents. As a result, the

use of cobalt is less critical. In their case, the element cobalt makes it possible to obtain a monozinc reagent from dihalo compounds of the same rank.

The electron-poverty of the nucleus may be

due either to the presence of electron-withdrawing
groups as substituents, or, in the case of six-membered
nuclei, to the replacement of a carbon with a hetero
atom. In other words, the electron-poor nucleus may be
a six-membered heterocyclic nucleus, especially
heterocyclic nuclei containing an atom from the
nitrogen column and more particularly nitrogen.

Among the electron-withdrawing groups that give good results, mention should be made of acyl groups, nitrile groups, sulfone groups, carboxylate

15 groups, trifluoromethyl groups or, more generally, perfluoroalkyl groups and halogens lower in rank than the halide which will be converted into an organozinc reagent. When the substituents are halogens of the same rank, a diorganozinc reagent is generally formed. These diorganozinc reagents constitute novel compounds and correspond to the general formulae below in which X and R both represent zinc-bearing groups.

Among the donor groups, that is to say groups giving mediocre results with chlorine but good results

25 with bromine, mention may be made of alkyloxy groups, alkyl groups, amine groups and dialkylamine groups.

The aromatic derivative which is the substrate in the present process advantageously corresponds to the following formula:



5 in which:

- Z represents a trivalent chain unit -C(R₁)=, an atom from column V, advantageously a nitrogen;
- X represents the leaving halogen;
- A represents either a chain unit chosen either from the groups ZH or from chalcogens advantageously at least equal in rank to that of sulfur, or from the two-membered divalent unsaturated groups CR₂=CR₃, N=CR₂, CR₂=N.

 $Insofar \ as \ they \ are \ borne \ by \ contiguous$ 15. atoms, two of the radicals R, R_1, R_2 and R_3 may be linked to form rings.

Thus, the aryls may especially be of formula:







in which ${\tt Z1}$ is chosen from the same meanings as those ${\tt 20}$ given for ${\tt Z.}$

 $\label{eq:R1R2} \mbox{The radicals R_1, R_2 and R_3 are chosen from the substituents mentioned above and especially:}$

 electron-withdrawing groups, in particular acyl groups, nitrile groups, sulfone groups, carboxylate

groups, trifluoromethyl groups or, more generally, perfluoroalkyl groups and halogens of a lower rank than the halide which will be converted into an organozinc reagent;

5 • donor groups, especially aryloxy or alkyloxy groups, hydrocarbyl groups such as aryls and alkyls (the latter word being taken in its etymological meaning) and amine groups, including groups monosubstituted and disubstituted with hydrocarbon-based alkylamine
10 groups.

It is desirable that the substrates should contain not more than 50 carbon atoms, advantageously not more than 30 carbon atoms and preferably not more than 20 carbon atoms.

Among the substrates that are particularly advantageous are halides, preferably arylchlorides, bearing, especially in the metaposition, an aliphatic carbon (that is to say an sp³ carbon) bearing at least two fluorines. For example, the halides are preferably trifluoromethylaryl chlorides.

This process for synthesizing organozinc reagents may be extended, firstly to all organozinc reagents linked to sp²-hybridized carbon atoms and especially to the synthesis of organozinc reagents from 25 vinyl halides, especially when these vinyl halides are conjugated with aromatic nuclei.

Although the technique is economically much less advantageous, it may also be advantageous to note that it can also be transposed to aliphatic halides.

One of the advantages of the present

5 invention is that it requires only complexing agents or ligands that are readily available, such as nitriles (preferably aromatic or bidentate) or pyridines and derivatives of the pyridine nucleus, such as quinoline.

Moreover, bipyridyls, being bidentate, also give good 10 results as ligands that are different from the solvent.

Although the bis-nitriles are capable of acting as bidentate ligands, they are poor complexing agents and should be used in high proportions of the same order as the monodentate ligands. They give good 15 results.

It is desirable, in order to avoid the medium being too acidic, that the bis-nitriles constituting the solvent, some of the solvent, or the ligand, should be such that, via the most direct pathway, two nitrile functions are separated by at least two carbons and advantageously three carbons.

Dinitroalkylenes in which the alkylene group contains from 2 to 8 carbon atoms give good results.

Mention may be made especially of glutaronitrile,

25 methylglutaronitrile, adiponitrile, pimelonitrile and suberonitrile.

25

Another advantage of the present invention is its ability to be performed readily at room temperature and, more generally, at a temperature below 50°C.

Finally, the reaction does not require an 5 inert electrolyte, since the zinc salt can be used as inert electrolyte.

Soluble zinc anodes may be used in this technique.

The present invention produces families of

organozinc compounds corresponding to the preceding
substrate formulae in which X has been replaced with a
zinc-bearing function (generally noted -Zn- X' in which
X' is halogen) which it has not been possible to obtain
previously. Among the families of interest which it has
not been possible to synthesize previously, mention
should be made of the compounds derived from the
preceding substrate formulae in which one of the
radicals R, R₁, R₂ and R₃ is a monosubstituted and
especially an unsubstituted aniline function.

Mention may also be made of compounds in which one of the radicals R, R_1 , R_2 and R_3 is a group bearing a sulfone group (-SO₂-), including sulfonates, that is vicinal to the aromatic nucleus, that is to say that it is adjacent to it.

Finally, the dizinc compounds in which R is a zinc-bearing group.

The characteristics of these families may be cumulated to form preferred subfamilies.

 $\label{eq:the_constraint} The \ non-limiting \ examples \ which \ follow \\ illustrate \ the \ invention.$

General procedure (condition A)

Apparatus

5 Single-compartment electrolysis cell equipped with a zinc anode and a nickel sponge cathode (gold or stainless steel cathodes may also especially be used).

Solvent: dimethylformamide/pyridine (45 ml/5 ml)

Ambient temperature (20 to 25°)

10 Aryl halide: 10 millimoles
Cobalt chloride: 1 millimole
Zinc bromide: 2.5 millimoles
Constant current: 0.2 A

No inert electrolyte

15 Electrode area: 20 cm²

Electrolysis time: 2 hours

The conditions differing from the general procedure are specified in the tables below, which give a sample of the results obtained.

20 The asterisk * indicates that the yield measurement was performed by coupling the organozinc reagent with phenyliodide.

5 with FG = electron-donating group

FGArX	FGArZnX %	ArH %	ArAr	Comments
PhCI	6	?	0	84% PhCl
PhBr	70	7	0	remaining
PhJ	20	Majority	0	All the PhI consumed
сн,о(С)Вг	82	18	0	
OCH ₃	75	21	o	
OCH,	62	37	0	
сн,о—О—а	6	20		74% ArCl remaining
сн,о-О-Вг	75	25	0	FG-Ar-Ph 70% Isolated*
N-O-Br	90	0	0	FG-Ar-Ph 85% Isolated*
H^N-\B1	85		0	

Table 2

In the case of aromatic halides

with FG = electron-withdrawing group

	WICH FG	- electio	ii-wir cua	rawing group
FGArX	ArZnX %	ArH	ArAr	Comments
MeO ₂ S————————————————————————————————————	90	ε	0	FG-Ar-Ph 80% Isolated
EsO ₂ C——Br	85			
COOEs	58			19% ArBr remaining
F ₃ C————————————————————————————————————	70			
Br—Br	70 only			
CI——Br	60 monozinc derivative (CIPhZnBr)			
F—Br	79			
CH2-C	25			disappearance of ArBr

 $\underline{ \mbox{Table 3}} \\ \mbox{In the case of heteroaromatic halides} \\$

ArX	ArZnX %
©N CI	50
N Br	25
S Br	67
S Br	25
Br—S—Br	20 (monozinc derivative)

5

In general, the thiophene derivatives show exceptional reactivity and it was possible in this case to perform a monoconversion of a dibromo derivative.

Example 4

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Table 4

Ir	the case of a	liphatic halid	es
RX	RZnX	RH	RY
	olo olo	ક <u>ુ</u>	
CH ₃ (CH ₂) ₃ Br	30	40	
Br (CH ₂) 3COOEt	42		*

Table 5

In	the case	of vinyl	. halide	
Ph-CH=CH	Br		PhCH=CHZnBr	45%

The tests below were carried out by varying

5 the operating conditions such as the nature of the anode, the concentration of catalyst, the concentration of zinc salts, or by using 2,2'-pyridine as ligand instead of pyridine.

10 Example 6

Table 6

ArX	ArZnX	Conditions
CHC——————————————————————————————————	75	$CoCl_2$, 1 eq, $ZnBr_2$, 2 eq, other conditions identical to A.
	50	CoCl ₂ 0.2 eq, other conditions identical to A.
	75	CoCl ₂ , 0.4 eq, other conditions identical to A.
	50	2 Bpy per Co, absence of pyridine, other conditions identical to A.
CH-C-CI	77	<pre>ZnBr₂, 2 eq, CoCl₂, 1 eq, other conditions identical to A</pre>
NC—()—Br	48	1 eq ZnBr ₂ , iron anode, other conditions identical to A.
	70	ZnBr ₂ , 2 eq, 1 eq CoCl ₂ , other conditions identical to A.
	70	ZnBr ₂ , 2 eq, 0.4 eq CoCl ₂ , other conditions identical to A.
ис-Сі	50	<pre>ZnBr₂, 2 eq, 0.4 eq CoCl₂, other conditions identical to A. ZnBr₂, 2 eq, CoCl₂, 1 eq, other</pre>
	65	conditions identical to A.

		22		
CN CI	60	60 ZnBr ₂ , 2 eq, CoCl ₂ , 0.4 eq, other conditions identical to A.		
Br	58	<pre>ZnBr₂, 2 eq, CoCl₂, 1 eq, other conditions identical to A.</pre>		
N Br	25	2 Bpy per CoCl ₂ , absence of pyridine, other conditions identical to A		

Example 7

Formation of organozinc reagents from ethyl parabromobenzoate, study of various solvents

THF made conductive with tetrabutylammonium fluoroborate gives good results for organozinc reagents, although slightly lower than in dimethylformamide. The other amides such as dimethylacetamide also give good yields of organozing 10 reagents. Nitriles such as acetonitrile give as much zinc reagent as when dimethylformamide is used.

Example 8

The results obtained below were achieved in 15 an acetonitrile/pyridine mixture (45/5). The other conditions were identical to the general conditions.

Table 7

Electrosynthesis of organozinc reagents in acetonitrile/pyridine medium (V/V = 9/1)

ArX	ArZnX
CF ₃ ,	80%
CH ₃ -C —————Br	57%
CH ₃ O ₂ S—Cl	90%
EtOOC Br	80%

The use of benzonitrile instead of pyridine (9/1 mixture by volume) also leads to good results under the general conditions. In particular, starting 5 with meta-bromofluorobenzene, a yield of 60% is obtained.

Example 9

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Ligands other than pyridine and benzonitrile
 Starting with pBr-PhCO₂Et under the general conditions described above (conditions A), the following results are obtained:

Solvent	Ligand	BrZnPhCO₂Et
Acetonitrile (45 ml)	Adiponitrile (5 ml)	75 %
Acetonitrile (45 ml)	1,2 - dimethoxyethane (5 ml)	65 %
acetonitrile (45 ml)	H₃C○N	50 %
,	(5 ml)	

2. Formation of aromatic and heteroaromatic dizinc reagents from aromatic dihalides (X-Ar-X)

General conditions identical to conditions A, but:

- the solvent is acetonitrile (45 ml)/pyridine
 (5 ml),
 - CoCl₂ 2 millimoles,
 - electrolysis stopped after 4 Faradays have been passed per mole of X-Ar-X (4 hours)

	23	
X-Ar-X	X-ArZn (%)	XZn-Ar-ZnX (%) 70
Br—O—Br	O	70
Br Br	Đ	51
Br Br	0	16
Br—CH ₃ —Br	0	66
Br-O-Br	0	65
CH ₃ O—O—Br	0	44
Br S Br	0	28
Br S Br	0	28
Cl—O—Br	38 Pcl-Ph-ZnBr	30
a-{O}-a	28	10
CI—CF ₃	73	19
a sta	26	2

CLAIMS

- The use of cobalt as a catalyst in the electrolytic synthesis of organozinc compounds, advantageously aryl or vinyl organozinc compounds.
- 5 2. The use as claimed in claim 1, characterized in that the cobalt is present in the electrolyte in oxidation state II.
- 3. The use as claimed in claims 1 and 2, characterized in that the cobalt is present in a 10 coordinated form.
 - 4. The use as claimed in claim 3, characterized in that the coordination of the cobalt is performed with a solutant or solvent compound that has a high donor number.
- 15 5. The use as claimed in claim 4, characterized in that the atom responsible for the good donor number is chosen from atoms of the nitrogen column.
 - The use as claimed in claims 3 to 5,
- 20 characterized in that the coordination of the cobalt is performed with a specific ligand.
- 7. The use as claimed in claims 3 to 6, characterized in that said ligand contains functions chosen from pyridine, nitrile, phosphine, stibine and 25 imine functions.

- A composition for electrolytic use, 8. -characterized in that it comprises a cobalt salt, a zinc salt, a solvent and a cobalt ligand.
- A process for the electrolytic synthesis 5 of organozinc reagents, advantageously aromatic or vinyl organozinc reagents, characterized in that it consists in subjecting a composition as claimed in claim 8, also comprising an organic halide, to an electrolysis on an inert cathode.
- 10. An aromatic organozinc compound comprising directly linked to an sp2 carbon atom, that is advantageously aromatic, at least one function or group chosen from not more than monosubstituted aniline functions, an SO2 group and another zinc-bearing 15 function.

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My residence, post office ad	dress and citizenship are as s	tated below next to my name.			
		e name is listed below) or an o d for which a patent is sought o			
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the specification of which (ch	neck only one item below):				
is attached her	eto.				
was filed as Ur	nited States application				
Serial No					
on					
and was amended					
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on June 30, 2000					
and amended under PCT ARTICLE 19					
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I acknowledge the duty to disclose information which is material to the patentability of this application in accordance with Title 37, Code of Federal Regulations. §1.56(a).					
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SIGNATURE OF INVENTOR 201	SIGNATURE OF INVENTOR 202	SIGNATURE OF INVENTOR 203
December 14 th, 2001	DATE December 14th 2001	DATE December 14th, 2001
	PAGE 2 OF 2	